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# High-throughput sequencing of transposable element insertions suggests adaptive evolution of the invasive Asian tiger mosquito towards temperate environments

Clement Goubert<sup>1,2,3,4</sup> | Helene Henri<sup>1,2,3</sup> | Guillaume Minard<sup>1,2,5,6</sup> | Claire Valiente Moro<sup>1,2,5</sup> | Patrick Mavingui<sup>1,2,5,7</sup> | Cristina Vieira<sup>1,2,3</sup> | Matthieu Boulesteix<sup>1,2,3</sup>

<sup>1</sup>Université de Lyon, Lyon, France

<sup>2</sup>Université Lyon 1, Villeurbanne, France

<sup>3</sup>Laboratoire de Biometrie et Biologie Evolutive, UMR CNRS 5558, Villeurbanne, France

<sup>4</sup>Department of Human Genetics, University of Utah, Salt Lake City, UT, USA

<sup>5</sup>Ecologie Microbienne, UMR CNRS 5557, UMR INRA 1418, Villeurbanne, France

<sup>6</sup>Department of Biosciences, Metapopulation Research Center, University of Helsinki, Helsinki, Finland

<sup>7</sup>UMR PIMIT, INSERM 1187, CNRS 9192, IRD 249, Plateforme Technologique CYROI, Université de La Reunion, Sainte-Clotilde, Reunion

## Correspondence

Matthieu Boulesteix, Laboratoire de Biometrie et Biologie Evolutive, UMR CNRS 5558, Université de Lyon 1, Villeurbanne, France  
Email: matthieu.boulesteix@univ-lyon1.fr

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## Abstract

Invasive species represent unique opportunities to evaluate the role of local adaptation during colonization of new environments. Among these species, the Asian tiger mosquito, *Aedes albopictus*, is a threatening vector of several human viral diseases, including dengue and chikungunya, and raises concerns about the Zika fever. Its broad presence in both temperate and tropical environments has been considered the reflection of great "ecological plasticity." However, no study has been conducted to assess the role of adaptive evolution in the ecological success of *Ae. albopictus* at the molecular level. In the present study, we performed a genomic scan to search for potential signatures of selection leading to local adaptation in one-hundred-forty field-collected mosquitoes from native populations of Vietnam and temperate invasive populations of Europe. High-throughput genotyping of transposable element insertions led to the discovery of more than 120,000 polymorphic loci, which, in their great majority, revealed a virtual absence of structure between the biogeographic areas. Nevertheless, 92 outlier loci showed a high level of differentiation between temperate and tropical populations. The majority of these loci segregate at high insertion frequencies among European populations, indicating that this pattern could have been caused by recent adaptive evolution events in temperate areas. An analysis of the overlapping and neighbouring genes highlighted several candidates, including diapause, lipid and juvenile hormone pathways.

## KEYWORDS

*Aedes albopictus*, genome scan, invasive species, local adaptation, transposable elements

## 1 | INTRODUCTION

Biological invasions represent unique opportunities to study rapid evolutionary changes, such as adaptive evolution. Settlement in a novel area is a biological challenge that invasive species have successfully overcome. The underlying processes can be studied at the molecular level, particularly to gather empirical knowledge of the genetics of invasions, a field of study that has produced extensive theoretical predictions for which there is still little evidence in nature (Colautti & Lau,

2015). Some of the main concerns are disentangling the effects of neutral processes during colonization, such as founder events or allele surfing at the migration front, from adaptive evolution (i.e., local adaptation, Colautti & Lau, 2015; Lande, 2015; Peischl & Excoffier, 2015).

Signatures of adaptation can be tracked on genomes due to characteristic patterns of reduced genetic diversity left by the appearance and spread of a new beneficial mutation (Braverman, Hudson, Kaplan, Langley, & Stephan, 1995; Fay & Wu, 2000;

Nielsen, 2005; Tajima, 1989; Vitti, Grossman, & Sabeti, 2013). Contrasting regimes of selection between populations can also leave high levels of genetic differentiation in the vicinity of adaptive loci (Lewontin & Krakauer, 1973; Maynard Smith & Haigh, 1974). The strength of such signals can be influenced by the origin of the adaptive mutation, for example, if it arises *de novo* or if it spreads from the standing genetic variation (Messer & Petrov, 2013; Pritchard, Pickrell, & Coop, 2010). However, detection of the footprint of natural selection is dependent on the availability of informative genetic markers, which should ideally provide substantial coverage of the genome to allow selection scans and be easily and confidently scored across many individuals. Unfortunately, invasive organisms are rarely model species, making the development of a reliable and efficient marker challenging.

The Asian tiger mosquito, *Aedes (Stegomyia) albopictus* (Diptera: Culicidae), is currently one of the most threatening invasive species (Invasive Species Specialist Group). Originating from southeastern Asia, it is one of the primary vectors of dengue and chikungunya viruses and is also involved in the transmission of other threatening arboviruses (Paupy, Delatte, Bagny, Corbel, & Fontenille, 2009), in particular, the newly emerging Zika virus (Chouin-Carreiro et al., 2016; Grard et al., 2014; Marcondes & Ximenes, 2015). Nowadays, *Ae. albopictus* has settled in every continent except Antarctica and is found in both tropical and temperate climates (Bonizzoni, Gasperi, Chen, & James, 2013). Although this species is supposed to have emerged from rain forests (Hawley, 1988), the acknowledged native area of *Ae. albopictus* encompasses contrasting environments including temperate regions of Japan and China, offering a large potential of fit towards the most recently colonized environments. For example, the induction of photoperiodic diapause in temperate areas, which has a genetic basis in *Ae. albopictus* (Hanson & Craig, 1994; Hawley, Reiter, Copeland, Pumpuni, & Craig, 1987; Urbanski, Benoit, Michaud, Denlinger, & Armbruster, 2010), is decisive to ensure invasive success in Europe or Northern America. It allows the susceptible populations to survive through winter at the larval stage into eggs. Such a trait appears to be governed by a “genetic toolkit” involving numerous genes and metabolic networks for which the genetic polymorphism between diapausing and nondiapausing strains remains to be elucidated (Poelchau, Reynolds, Elsik, Denlinger, & Armbruster, 2013a). In addition, the colonization of new areas that appear similar to the native environment at first glance can still involve *de novo* adaptation; even environments that share climatic variables are not necessarily similar regarding edaphic and biotic interactions (Colautti & Lau, 2015). This suggests that, regardless of the native and settled environments, it might be possible to find evidence of adaptive evolution in invasive populations of *Ae. albopictus*.

To better understand the invasive success of this species, we genotyped 140 field individuals collected from three Vietnamese (native tropical area) and five European (invasive temperate area) populations (Figure 1), aiming to identify genomic regions involved in local adaptation. To do so, we developed new genetic markers based on high-throughput genotyping of the insertion of transposable

elements (TEs) that represent at least one-third of the genome of *Ae. albopictus* and include recently active families that can reach thousands of copies in one genome (Goubert et al., 2015).

Amplification of TE insertions is particularly efficient to obtain many genetic markers throughout one genome (Bonin et al., 2008), especially if few genomic resources are available (Monden, Yamaguchi, & Tahara, 2014), which was the case for the Asian tiger mosquito until recently. In addition, such markers represent an attractive alternative to other methods of diversity reduction such as RAD sequencing (Miller, Dunham, Amores, Cresko, & Johnson, 2007) that could be less efficient in species with a high TE load (Davey et al., 2013) and did not produce satisfying results in *Ae. albopictus* (Goubert, Minard, Vieira, & Boulesteix, 2016). In mosquitoes, TEs have been shown to be powerful markers for both population structure analysis (Biedler et al., 2003; Boulesteix et al., 2007; Esnault et al., 2008; Santolamazza et al., 2008) and genome scans (Bonin, Paris, Tetreau, David, & Després, 2009).

We hypothesized that some TE insertion sites could be located in the neighbourhood of targets of natural selection and thus could reach a high level of differentiation between native and invasive populations if selective sweeps occurred during local adaptation. In addition, some TEs could also insert near or inside coding regions and many studies have shown their recurrent involvement in environmental adaptation in multiple organisms (Casacuberta & González, 2013), eventually contributing to the success of invasive species (Schrader et al., 2014; Stapley, Santure, & Dennis, 2015).

To distinguish between neutral demographic effects and adaptive evolution, we first performed population genetic analyses to reveal the global genetic structure of the studied populations. We then performed a genomic scan for selection and identified 92 candidate loci under directional selection, among which several are located within or in a close neighbourhood of annotated genes, revealing candidate pathways to investigate in forthcoming studies.

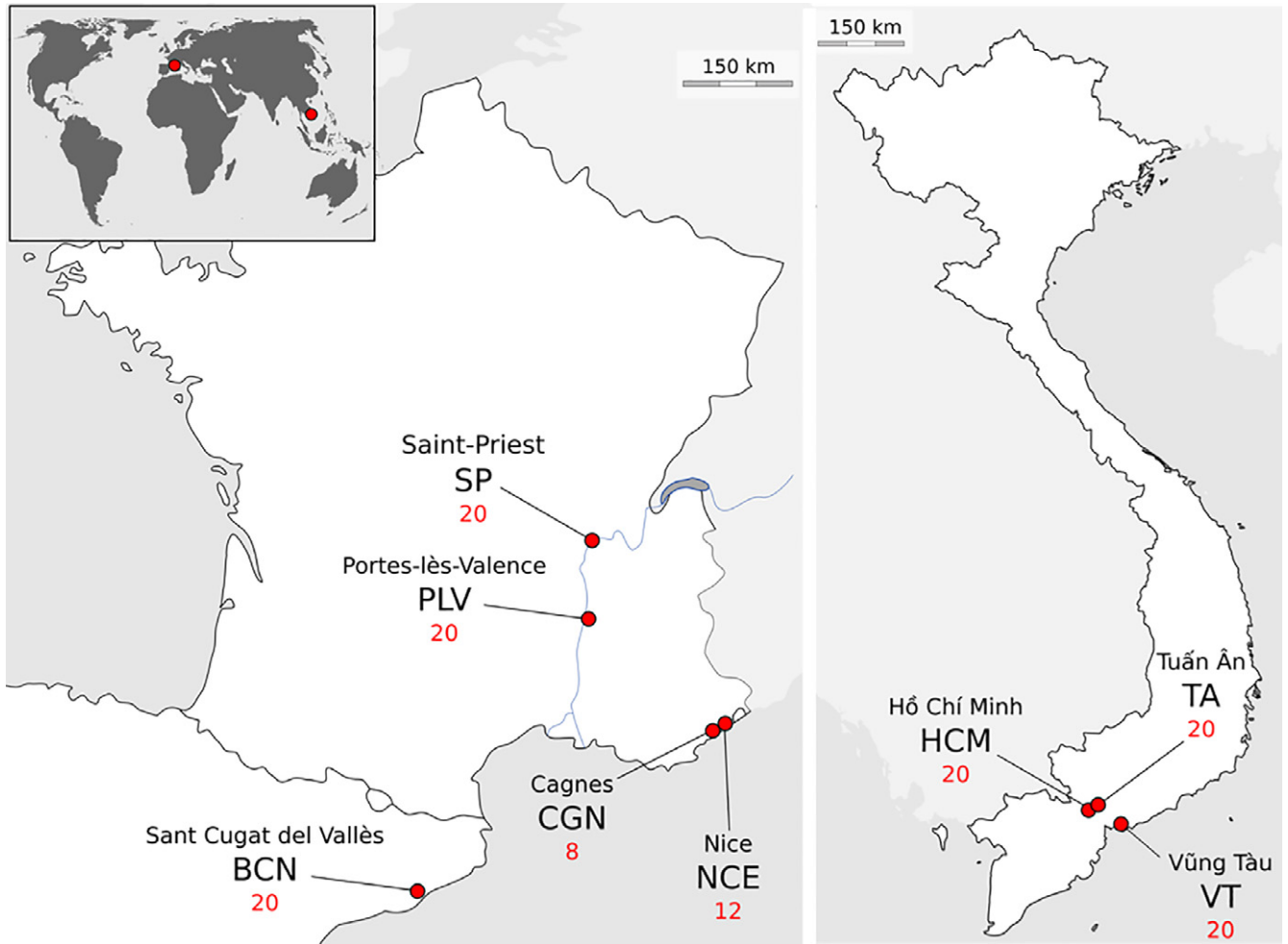
## 2 | MATERIALS AND METHODS

### 2.1 | Biological samples

One hundred and forty flying adult female *Ae. albopictus* were collected in the field at eight sampling sites in Europe and Vietnam during the summers of 2012 and 2013 (Figure 1 and Table S1). Individuals were sampled using either a single trap or aspirators through the sampling site within a 50 m radius. When traps were used, live mosquitoes were collected after a maximum of 2 days.

### 2.2 | High-throughput transposon display (TD) genotyping

The insertion polymorphism of the five transposable element families (I Loner Ele1 (IL1), Loa Ele2B (L2B), RTE4, RTE5 and Lian 1) identified by Goubert et al. (2015) in *Ae. albopictus* was characterized using transposon display (TD), a TE insertion-specific PCR method, combined with Illumina sequencing of all TD amplification products



**FIGURE 1** Sampling sites (with abbreviations) of *Ae. albopictus* in Europe and Vietnam. Red numbers correspond to the total number of individuals sampled. Supporting information on samples is available in Table S1

(Fig. S1). These TE families were chosen according to high copy number estimate (from 513 to 4,203 copies), high identity between copies and a “copy and paste” mode of transposition (all these TEs are non-LTR retrotransposons).

### 2.2.1 | DNA extraction and TD adapter ligation

The total DNA was extracted from whole adult bodies following the phenol-chloroform protocol described by Minard et al. (2015). The TD was conducted combining methods from previous studies (Akkouche et al., 2012; Carnelossi et al., 2014; Munroe et al., 1994; Roy et al., 1999). First, individual extracted DNA ( $\approx 75$  ng) was used for enzymatic digestion in a total volume of 20  $\mu$ l, with HindIII enzyme (10 U/ $\mu$ l) and buffer R (Thermo Scientific) for 3 hr at 37°C. The enzyme was inactivated at 80°C for 20 min. TD adapters were then built by hybridizing Hindlink with MSEB oligonucleotides (100  $\mu$ M, see Table S2) in 20 $\times$  SSC and 1 M Tris in a total volume of 333  $\mu$ l after 5 min of initial denaturation at 92°C and 1 h at room temperature for hybridization. Once ready, the TD adapters were ligated to 20  $\mu$ l of the digested DNA by mixing 2  $\mu$ l of TD adapter

with 10 U T4 ligase and 5 $\times$  buffer (Fermentas) in a final volume of 50  $\mu$ l for 3 hr at 23°C.

### 2.2.2 | Library construction

For each individual and for each of the five TE families, the TE insertions were amplified by PCR (PCR 1) in a Bio-Rad Thermal Cycler (C1000 or S1000) in a final volume of 25  $\mu$ l. The mixture contained 2  $\mu$ l of digested-ligated DNA with 1  $\mu$ l dNTPs (10 mM), 0.5  $\mu$ l TD adapter-specific primer (LNP, 10  $\mu$ M, see Table S3) and 0.5  $\mu$ l of TE-specific primer (10  $\mu$ M), 1 U AccuTaq polymerase (5 U/ $\mu$ l) with 10 $\times$  buffer and dimethyl sulphoxide (Sigma). Amplification was performed as follows: denaturation at 98°C for 30 s then 30 cycles of 94°C for 15 s, hybridization at 60°C for 20 s and elongation at 68°C for 1 min; a final elongation was performed for 5 min at 68°C. For L2B and RTE5 TEs, a nested PCR was performed to increase specificity under the same PCR conditions using internal forward TE primers and LNP (Table S2). The PCR 1 primers include a shared tag sequence that was used for hybridization of the individual indexes by PCR 2.

Multiple independent PCR 1 can be performed to avoid amplification bias during the library preparation (Recknagel, Jacobs, Herzyk, & Elmer, 2015). Accordingly, three independent PCR 1 were performed from the same digestion product for each TE family. The PCR 1 products (3 PCR \* 5 TE per individual) were then purified using volume-to-volume Agencourt AMPure XP beads (20  $\mu$ l PCR 1 + 20  $\mu$ l beads) and eluted in 30  $\mu$ l resuspension buffer. After NanoDrop quantification, equimolar pools containing the 3\*5 PCR products per individual were made using a Tecan EVO200 robot. Individual pools were then size selected for 300- to 600-bp fragments using Agencourt AMPure XP beads as follows: first, the magnetic beads were diluted in H<sub>2</sub>O at a 1:0.68 ratio then added to 0.625 $\times$  PCR products to exclude long fragments. A second purification was performed using a nondiluted bead:DNA ratio of 1:8.3 to exclude small fragments.

Multiplexing samples was performed using a homemade 6 bp index (included in SRA individual name), which was added to the R primer (Table S2) during a second PCR (PCR 2) with 12 cycles in an ABI 2720 Thermal Cycler. The mixture contained 15 ng PCR products, 1  $\mu$ l of dNTPs (10 mM), 0.5  $\mu$ l MTP Taq DNA polymerase (5 U/ $\mu$ l, Sigma), 5  $\mu$ l 10 $\times$  MTP Taq buffer and 1.25  $\mu$ l of each tagged primer (20  $\mu$ M) in a final volume of 50  $\mu$ l. Amplification was performed as follows: denaturation at 94°C for 60 s and then 12 cycles of denaturation at 94°C for 60 s, hybridization at 65°C for 60 s and elongation at 72°C for 60 s; a final elongation was performed for 10 min at 72°C. The PCR 2 products were purified using an Agencourt AMPure XP bead:DNA ratio of 1:1.25 to obtain libraries. TD product purification, library preparation and paired-end sequencing using an Illumina HiSeq 2000 (1 lane) was performed at the GeT-PlaGe core facility (Genome and Transcriptome, Toulouse) using a TRUSEQ PE CLUSTER KIT version 3 (2  $\times$  100 bp) and a TRUSEQ SBS KIT version 3.

## 2.3 | Bioinformatic treatment of TD sequencing

The steps of the informatics treatment from the raw sequencing data set to population binary (1/0) matrices for the presence/absence of TE insertions per individual are described in Fig. S2. First (Fig. S2-A), the paired-end reads of each individual were quality-checked and trimmed using *URQT* version 1.0.17 (Modolo & Lerat, 2015) with standard parameters and a *t* quality threshold of 10. The reads pairs were then checked and trimmed for Illumina adapter contamination using *Cutadapt* (Martin, 2011). The R1 reads include the TE sequence and the R2 reads include the TD adapters followed by the flanking sequence of the closest *EcoRI* site, called here the “insertion locus” (Fig. S2A). Specific amplification of TE insertions was controlled by checking for the expected 3' TE sequence on the R1 read using *Blat* (Kent, 2002) with an identity threshold of 0.90. Only reads with an alignment-length/read-length ratio  $\geq 0.90$  were retained. The R2 reads for which the R1 mate passed this filter were then selected for insertion loci construction after removal of the TD adapter on the 5' start using *Cutadapt* and removal of reads under 30 bp. Selected reads were separated in each individual according to the TE families for loci construction.

To correct for the interindividual coverage variations, we performed a sampling of the cleaned reads (Fig. S2-B). First, for each TE family, the distribution of the number of reads per individual was generated and individuals with fewer reads than the first decile of the distribution were removed; then, the cleaned reads of the remaining individuals were randomly sampled at the value of the first decile of coverage (this value varies among TE families). For each TE, the sampled reads of each retained individual were clustered together using the *CD-HIT-EST* program (Li & Godzik, 2006) to recover insertion loci (Fig. S2-C). In this all-to-all reads comparison, the alignments needed a minimum of 90% identity, the shortest sequence needed to be 95% the length of the longest, global identity was used and each read was assigned to its best cluster. In a second step, the reference reads of each locus within an individual, given by *CD-HIT-EST*, were clustered with all reference reads of all individuals using the same threshold to build the locus catalog, including a list of loci of all individuals and the coverage for each locus in each individual. After this step, the insertion loci that matched known repeats of the Asian tiger mosquito (Goubert et al., 2015) were discarded; alignments were performed with *Blastn* using the default parameters.

As the quality control removed a substantial number of reads for the construction of the TE insertions catalog, the raw R2 reads (with their TD adapter removed), which could have been discarded in the first attempt, were then mapped over the catalog to increase the scoring sensibility (Fig. S2-D). Before mapping, the raw R2 reads were also sampled at the first decile of individual coverage (as described previously). At this step, individuals who were removed from at least two TE families for loci construction were definitively removed from the whole analysis. Mapping (Fig. S2-E) was performed over all the insertion loci of all TE families in a single run to prevent multiple hits. *Blat* was used with an identity threshold of 90%. Visual inspection of alignment quality over 30 sampled loci per TE family was performed to ensure the quality of scoring. Raw matrices were then filtered out (Fig. S2-F) for a minimum insertion frequency of 2.5% among all individuals and aberrant loci with extreme ( $>99$ th centile) coverage and coverage standard deviations were discarded. The final data sets consisted of one matrix per TE family with information for each individual concerning the presence (1) or absence (0) of TE for each of the selected loci.

To check if the sampling procedure would affect our results, the read sampling procedures and subsequent analysis were performed independently 3 times (replicates M1, M2 and M3).

## 2.4 | Genetic analyses and genomic scan

The population structure analyses were performed independently for each TE family. Principal coordinates analyses (PCoAs) were performed to identify genetic clusters using the *ADE4* package (Dray & Dufour, 2007) of *R* 3.2.1 (R Core Team 2016). The *S7* coefficient of Gower and Legendre (1986) was used as a genetic distance because it gives more weight to shared insertions as follows: with *a*, *b*, *c* and *d* taken from a contingency table such as *a* = 1/1 (shared presence); *b* = 1/0; *c* = 0/1; and *d* = 0/0 (shared absence);  $S7 = 2a/(2a+b+c)$ .



Shared absences were not used because they do not provide information on the genetic distance between individuals due to the “copy and paste” mode of transposition of the TE used (shared ancestral state). In order to obtain a quantitative measure of the genetic differentiation, pairwise  $F_{ST}$  between populations were computed using ARLEQUIN 3.5 (Excoffier & Lischer, 2010); the significance of the index was assessed over 1,000 permutations using a significance threshold of 0.05.

The genomic scan was performed in two steps for each of the sampling replicates of each TE. First, BAYESCAN 2.1 (Foll & Gaggiotti, 2008) was used to test for the deviation of each locus from neutrality. BAYESCAN considers a fission/island model in which all subpopulations are derived from a unique ancestral population. In this model, variance in allele frequencies between subpopulations is expected to be due either to the genetic drift that occurred independently in each subpopulation or to selection that is a locus-specific parameter. The differentiation at each locus in each subpopulation from the ancestral population is thus decomposed into a  $\beta$  component (shared by all loci in a subpopulation) related to genetic drift and a  $\alpha$  component (shared at a locus by all subpopulations) due to selection. Using a Bayesian framework, BAYESCAN tests for the significance of the  $\alpha$  component at each locus. Rejection of the neutral model at one locus is conducted using posterior Bayesian probabilities and controlled for multiple testing using a false discovery rate. In addition, BAYESCAN integrates uncertainty about allele frequency from dominant data such as the TD polymorphism, leaving the inbreeding coefficient ( $F_{IS}$ ) to vary between 0 and 1 during the Markov chain Monte Carlo process. BAYESCAN was used with default values except for the prior odds, which were set to 100 (more compatible with data sets having thousands of loci, see BAYESCAN manual) and a significance  $q$ -value threshold of 0.05 to retain outlier loci.

In a second step, only outliers suggesting divergent directional selection between Europe and Vietnam were considered. To identify these, locus-by-locus analyses of molecular variance (AMOVAs) were performed using ARLEQUIN 3.5 for each TE family. The significance of the  $F_{CT}$  (hierarchical analogue of the  $F_{ST}$  measuring the extent of differentiation between groups of populations) between Vietnamese and European populations was assessed by performing 10,000 permutations between individuals among populations with a significance threshold of 0.05. For each data set, BAYESCAN outliers were cross-referenced with significant  $F_{CT}$  loci as an objective threshold to restrict the number of candidate loci.

## 2.5 | PCR validation and Outlier analyses

Pairs of primers were designed for each outlier locus to be used in standardized conditions. The forward primer was located in the TE end of the concerned family, and the reverse primer was set from the outlier locus (Table S2). All primer pairs were first tested on a set of 10 individuals to assess their specificity using a 1/50 dilution of starting DNA from the TD experiment. Validated primers were then used to check the insertion polymorphisms in 47 representative individuals from the eight populations studied in the TD experiment

using 1/50 dilutions of the starting DNA (not all individuals could be used because of DNA limitations). All PCRs were conducted in a final volume of 25  $\mu$ l using 0.5  $\mu$ l of diluted DNA, 0.5  $\mu$ l of each primer (10  $\mu$ M), 1  $\mu$ l of dNTPs (10 mM) and 1 U of DreamTaq polymerase with 1 $\times$  green buffer (ThermoFisher Scientific). Amplification was performed as follows: denaturation at 94°C for 2 min and then 34 cycles including denaturation at 94°C for 30 s, hybridization at 60°C for 45 s and elongation at 72°C for 45 s; a final elongation was performed for 10 min at 72°C. After a 45-min migration of the PCR product on a 1 $\times$  electrophoresis agarose gel, CG and MB assessed the insertion polymorphisms independently.

To identify the genomic environment of the outlier loci, their sequences (reference R2 read) were mapped onto the assembled genome of *Ae. albopictus* assembly AaloF1 strain Foshan (Chen et al., 2015) using Blastn. Blastn alignments were performed with default parameters and sorted according to their score. Additionally, each alignment was visually inspected for consistency. Outlier loci with multiple identical hits were discarded. To identify genes surrounding the mapped outliers, we used annotations from VectorBase (<http://www.vectorbase.org>) for the assembly or, if the *Ae. albopictus* gene was missing annotation, we used the annotation of orthologues in closely related species. In addition, we questioned whether genes potentially involved in the diapause pathway, a critical adaptation required in temperate environments, may be associated with outliers. In a previous publication, Poelchau et al. (2013a) identified that differentially expressed genes between diapause-induced and nondiapause-induced samples were significantly enriched in functional categories related to diapause preparation. These functional categories are defined a priori and tested for a more significant differential expression than that expected by chance (Poelchau et al., 2013a). Because, at the time of this previous publication, the corresponding transcripts were associated with *Ae. aegypti* orthologues, we here mapped the original transcriptome data (annotated eggs and embryo assembly, downloaded at <http://www.albopictusexpression.org>) onto the AaloF1 genome assembly using Blat with default parameters. After alignment, one best hit was retained per transcript according to the best alignment score. When a transcript had identical best hits, all positions for the transcript were considered. After alignments, the transcript positions were intersected with the AaloF1 gene set using BEDTOOLS version 2.25.0 (Quinlan & Hall, 2010) to identify corresponding *Ae. albopictus* genes. In addition, Chen et al. (2015) reported 71 complete diapause-related genes that were merged to our initial candidate set. The eventual enrichment in outliers near these diapause-related genes was assessed using the following procedure: we first estimated the total number of base pairs covered by diapause-related genes on the genome assembly. We then added a fixed distance in 5' and 3' of each of these genes (up to the contig size) corresponding to the longest distance observed between gene and outlier among the outliers found close to diapause-related gene; this defined the “diapause base pairs.” We then compared the ratio of the number of outliers found within these “diapause base pairs” over the total number of mapped outliers and the ratio of “diapause base pairs” over the total assembly size using an exact binomial test.

### 3 | RESULTS

#### 3.1 | High-throughput TE insertion genotyping

The presence/absence of insertions of five TE families was genotyped in an initial number of 140 individuals using a combination of family-specific PCR and individually labelled high-throughput sequencing. Sequencing produced a total of 102,319,300 paired-end reads ( $2 \times 100$  bp). After quality and specificity filtering, 24,332,715 reads were suitable for analyses. The loss of reads was in the great majority due to specificity filtering as quality only resulted in trimming. After application of the read sampling procedure to control for coverage variation between individuals, an average of 128,491 polymorphic insertion loci was available for each of the three sampling replicates. A final 120 individuals were retained per TE family (discarded individuals vary per TE family). The mean number of loci per individual and per TE family ranged from  $1,025 \pm 290$  SD (IL1 family, mean and SD averaged over the three replicates) to  $3,266 \pm 766$  SD (RTE5 family). Details are given in Table S1. Although our read sampling procedure could have artificially lowered the mean insertion frequency of the loci, this effect should be small because in our final data sets the TE insertion frequencies (i.e., the number of individuals who share an insertion) are not influenced by the mean number of reads per individual at the considered locus (Fig. S3). Additionally, we did not detect any bias between European and Vietnamese populations when measuring the identity between the R1 reads (TE part) and the TE consensus sequence of each TE family, suggesting that the relative activity of the TE is comparable between continents (Fig. S4).

#### 3.2 | Population structure

Principal coordinates analyses (PcoAs) were performed independently for each of the five TEs (Figure 2). The shared absence of a specific insertion was not considered in the distance matrices in the PCoAs: Class I retrotransposons have a “copy and paste” transposition mechanism that allows us to infer the “absence” state as the ancestral allele. In addition, these TE insertions segregate at very low frequencies among individuals and thus a shared “absence” is likely to be noninformative with regard to an individual's co-ancestry. Among the three main principal coordinates (PCs), individuals tend to be grouped according to their respective populations with little overlap between groups. However, the three main PCs represent only a small fraction of the total variance (<10%), suggesting a weak genetic structuring between the populations. Overall, individuals from Vietnamese populations (HCM, TA, VT) tend to be grouped together in a single cluster, with the exception of 13 to 14 individuals from HCM for the L2B and RTE5 TE families (Fig. S5) and six individuals of VT with the RTE4 TE family (Figure 1) that cannot be clearly distinguished from European samples. BCN individuals (Spain) represent the most homogeneous group, well differentiated from Vietnamese and French individuals (SP, CGN, NCE and PLV).

In agreement with the PCoAs, the analyses of molecular variance (AMOVAs) attributed little genetic variance among groups

(Vietnam–Europe) and between populations within groups (Table 1). In the studied populations, most of the genetic variance was distributed among individuals within groups.

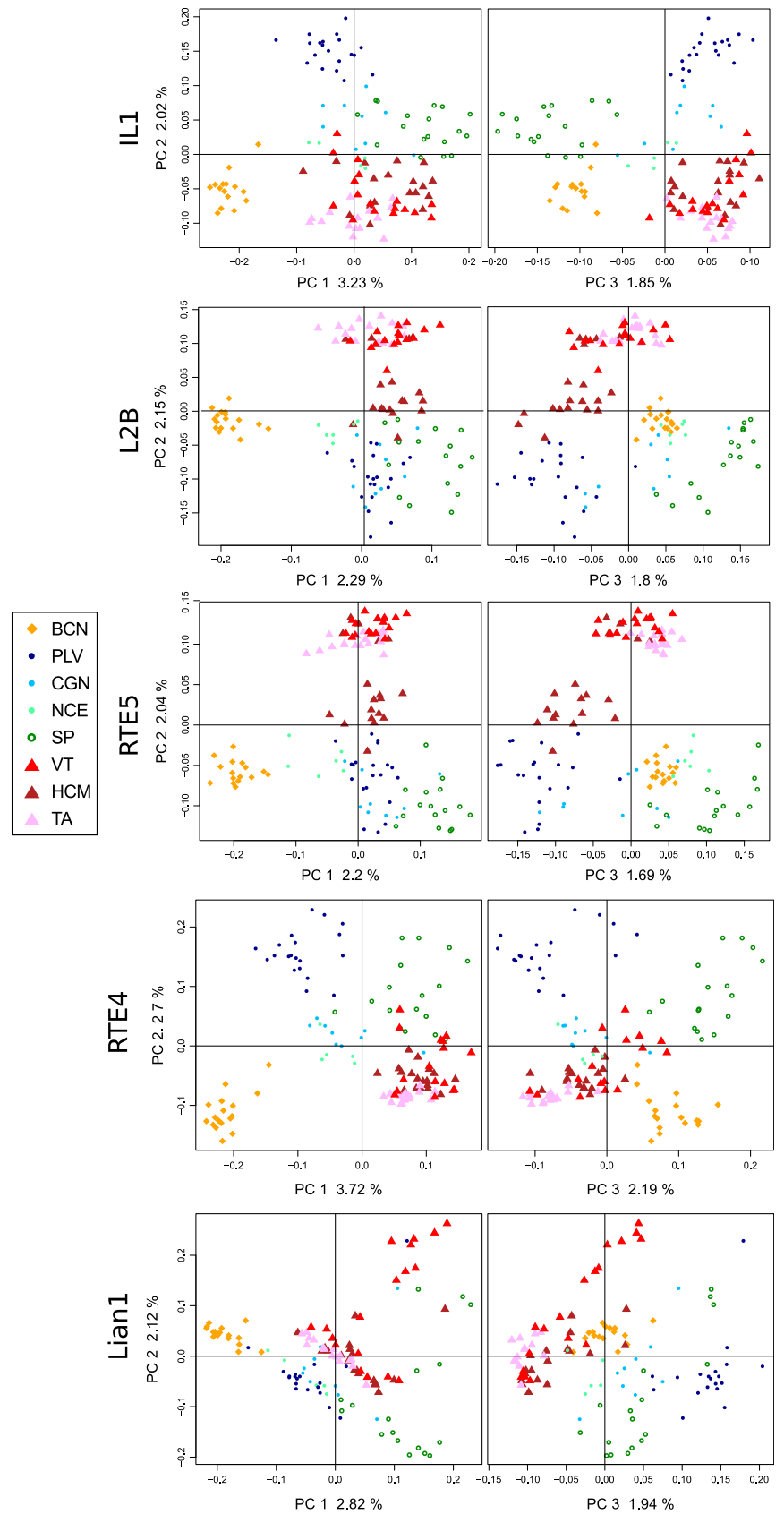
The measures of genetic differentiation among pairs of populations were consistent between the PCoAs and AMOVAs (File S1): the BCN population shows the highest  $F_{ST}$  with the other populations for each of the five TEs ( $0.051 < F_{ST} < 0.148$ ), whereas Vietnamese populations were the most closely related ( $0.011 < F_{ST} < 0.032$ ). Although VT is located 100 km away from TA and HCM (both sampled in the same city, Hô Chi Minh, Vietnam), the  $F_{ST}$  values are very similar between the three Vietnamese populations, suggesting no influence of geography at this scale. CGN and NCE, sampled in the same urban area (Nice agglomeration), also differentiated little or were not significantly differentiated, depending on the TE family. The previously identified intermediate pattern of HCM with some European populations at the L2B and RTE5 loci (PCoAs) is also found at the  $F_{ST}$  level, especially for the low differentiation with the PLV population for these markers ( $0.011 < F_{ST} < 0.020$ ). Neighbor-joining trees of individuals based on the S7 coefficient of Gower and Legendre (1986) also confirmed the weak population structure of the sampled population of *Ae. albopictus* (Fig. S6) and are characterized by especially long branches due to a high amount of low frequency insertions.

#### 3.3 | Genomic scan

Outlier loci for selection signature were searched using BAYESCAN (nonhierarchical island model) and then sorted for a significant  $F_{CT}$  (between Europe–Vietnam group differentiation) to retain only candidate loci compatible with a differential selection between continents. To reduce false-positive risks due to uneven mutation rates between TE families, outlier scans were also performed independently for each TE family (Narum & Hess, 2011; de Villemereuil, Frichot, Bazin, François, & Gaggiotti, 2014). We identified 92 candidate insertion loci (Figures 3, S7 and Table S4). Most of these insertions are found in both areas (no private allele), except for RTE4\_1638 and RTE4\_1898 that were not found in Vietnam. In addition, 74% of the outliers correspond to high-frequency insertions in Europe, which is significantly more than expected for a 50-50 chance (chi-squared test,  $X = 20.098$ ,  $p < .01$ ), whereas this 50-50 pattern is observed comparing 92 randomly chosen loci with the same overall insertion frequency ( $\geq 20$  individuals/locus, chi-squared test,  $X = 1.837$ ,  $p = .175$ ) between Europe and Vietnam (Figure 4).

PCR amplification of the outlier loci was carried out on a representative panel of 47 individuals to validate the insertion pattern detected using TD (see Materials and Methods). For loci where the amplification was successful, the insertion pattern observed using PCR always confirmed the TD pattern (Fig. S8).

From 92 outlier loci, 21 could be attributed to a unique position on the *Ae. albopictus* genome. Annotation and distance to surrounding genes are reported in the File S2. Fifteen outliers mapped within contigs with identified genes. We found that four outliers (File S2, sheet 2, highlighted) loci are located on contigs that harbour



**FIGURE 2** Principal coordinates analyses (PCoAs). Projection of individuals over the three-first principal coordinates (PC) of PCoAs for each of the five TE families and for the first replicate (M1, see Materials and Methods). The proportion of inertia represented by each axes is noted in %. Circles: European populations; triangles: Vietnamese populations. The results for other sampling replicates can be found in Figure S5

diapause-related genes. Two of them (Lian1\_5902 and RTE4\_17015) are located in the direct vicinity, either inside or within 5,654 bp, of these genes, which is significantly closer than expected by chance

(exact binomial test,  $p = .014$ ). Lian1\_5902 is located in an intron of *lac1* (longevity assurance factor 1; AALF000670), and RTE4\_17015 neighbours the AALF004790 a lipophorin-coding gene. Both genes



**TABLE 1** Analyses of molecular variance (AMOVAs). The results for the three replicates (M1, M2, M3) of read sampling for the five TE families (IL1, L2B, RTE5, RTE4, Lian1). Values are given in percentage of the total genetic variance

	IL1	L2B	RTE5	RTE4	Lian1
Among groups	[0.59–0.70]	[1.22–1.29]	[1.08–1.10]	[1.97–2.04]	[0.67–0.74]
Among populations within groups	[5.15–5.37]	[3.58–3.63]	[3.36–3.40]	[6.67–6.78]	[4.47–4.56]
Within populations	[94.04–94.16]	[95.08–95.18]	[95.51–95.55]	[91.18–91.30]	[94.77–94.81]

Intervals report min and max values among the three sampling replicates.

are known to be involved in lipid metabolic pathways. Although the other diapause-related genes are not the closest genes of the two other outliers, they represent two singular groups of genes located in tandem: AALF020842 and AALF00843 located 71.05 kb from RTE5\_10123 and AALF020959, AALF020960, AALF020961, AALF020962, AALF020963 and AALF020965 located 216.6 kb from Lian1\_11252.

Three other outliers are located within other genes (outliers Lian1\_10005, Lian1\_9293 and RTE4\_34941), including a hemolymph juvenile hormone-binding protein (RTE4\_34941/AALF012643). The 10 remaining outliers were located 21.1–85.2 kb from their closest gene.

## 4 | DISCUSSION

The goal of our study was to identify genomic regions involved in adaptive evolution of *Ae. albopictus* thanks to the development of new genetic markers. Through high-throughput genotyping of the insertion polymorphisms of five TE families, we identified up to 128,617 polymorphic loci among more than a hundred of individuals from eight sampling sites. The estimated genome size of *Ae. albopictus* exceeds one billion base pairs (Chen et al., 2015; Dritsou et al., 2015; Goubert et al., 2015). Accordingly, the number of markers scored in this study offers a comfortable genomic density of one marker every 10 kb.

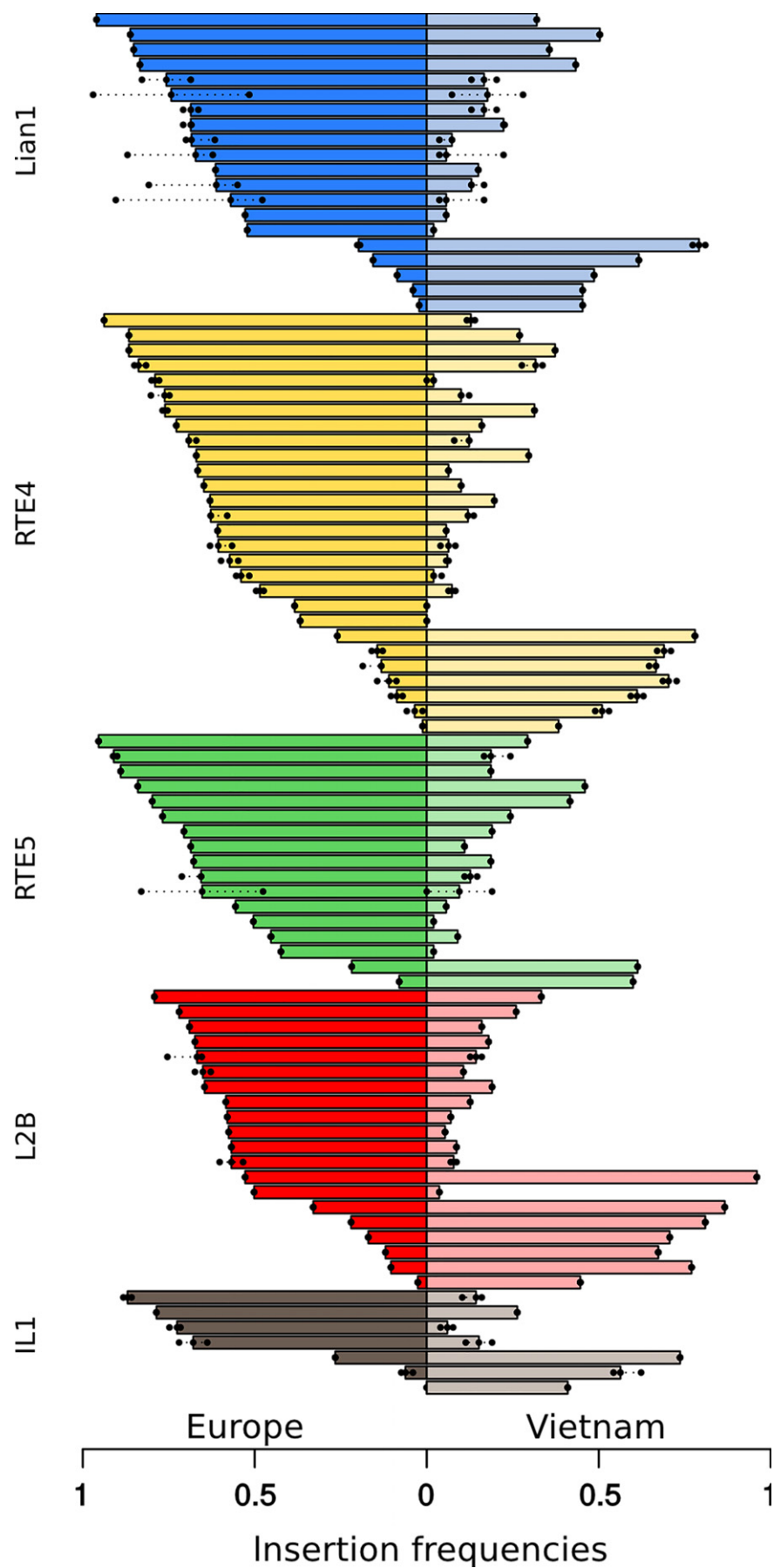
TE-based methods have been successfully used to perform population genetic analyses within a repetitive genomic environment, such as in the human genome (Rishishwar et al., 2015; Watkins et al., 2003; Witherspoon et al., 2013). Similar high-throughput genotyping methods have been developed for a large panel of organisms (Bridier-Nahmias et al., 2015; Iskow et al., 2010; Monden & Tahara, 2015; Sabot et al., 2011; Witherspoon et al., 2010) but relied on well-established reference genomes (human, rice, strawberry, yeast). Monden et al. (2014) recently completed such an analysis without a reference genome, to score 2,024 loci from two TE families in sweet potato. Because of the number of available loci, and being the first of its kind in animals, our study represents a large improvement. We provide a cost-efficient method to generate many polymorphic markers without extensive knowledge of a species' genome. Specifically, this strategy appears valuable for species with a large genome size in which the TE density could severely compromise the development of more classical approaches.

Genotyping from high-throughput sequencing keeps uncertain whether the observed absence of a TE insertion is true or is due to

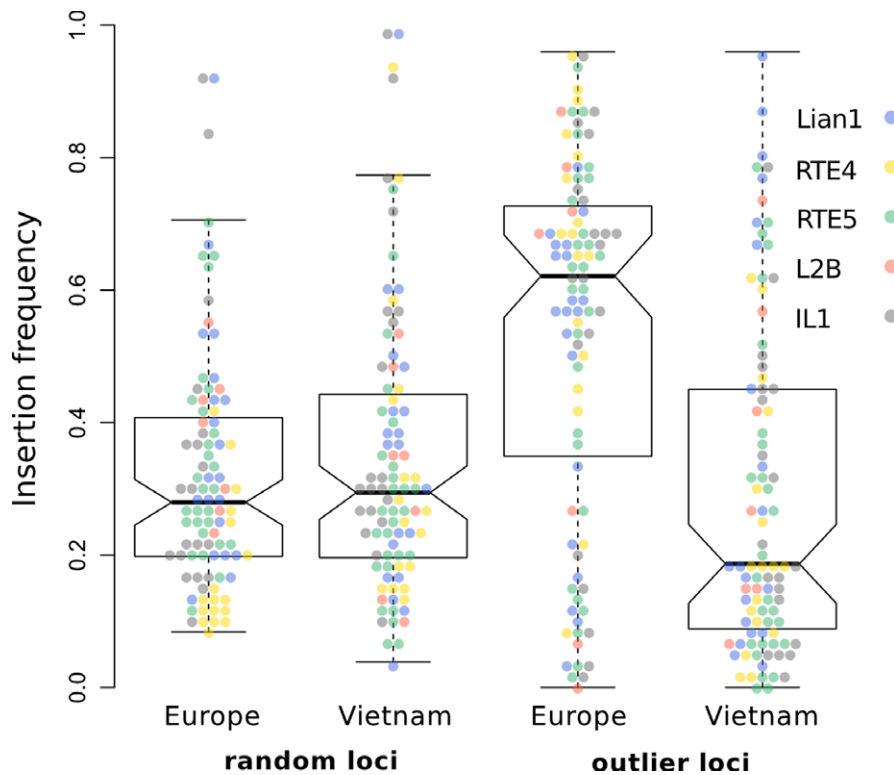
a lack of coverage. However, we showed that there is no correlation between the TE insertion frequencies and the mean coverage of loci (Fig. S3), which should be the case if false negatives were systematically associated with a lack of coverage. Moreover, the genetic structure of the studied populations showed strong consistency between sampling replicates of individual's reads (Fig. S5), demonstrating the robustness of the method despite an initial substantial coverage variation among individuals. Population genetics analyses revealed a very low level of genetic structuring between European and Vietnamese populations. Among the studied populations, AMOVAs showed that most of the genetic variation is distributed between individuals within populations (>90%), and as suggested by pairwise  $F_{ST}$  and PCoAs, only a small part (<10%) of the genetic variance is due to differentiation between populations. The genetic differentiation we measured is as high among European populations as it is between populations from Europe and Vietnam.

This singular population structure is in agreement with previous results gathered in *Ae. albopictus* using different collections of allozymes, mtDNA or microsatellite markers (Black, Ferrari, & Sprengert, 1988; Gupta & Preet, 2014; Kambhampati, Black, & Rai, 1991; Manni et al., 2015; Zhong et al., 2013). Moreover, a recent analysis performed with a set of 11 microsatellites on individuals from the same populations (with the exception of BCN) showed a similar distribution of genetic variation among hierarchical levels (Minard et al., 2015). These results demonstrate the reliability of our markers and confirm that a nonhierarchical island model can likely fit the global genetic structure. The genetic diversity observed in Europe is compatible with a scenario of multiple and independent introductions, as already suggested for *Ae. albopictus* (Becker et al., 2013; Birungi & Munstermann, 2002; Takumi et al., 2009; Urbanelli, Bellini, Carrieri, Sallicandro, & Celli, 2000). However, as previously suggested, this pattern could also be the result of founder events that may occur during colonization and/or a restriction of gene flow between populations after their introduction. Answering such a question would require extended sampling over the entire native area.

The outlier analysis revealed 92 loci with high posterior probabilities of being under positive selection between European and Vietnamese populations. When possible, the PCR amplification of the outlier loci using a set of representative individuals confirmed a shift of insertion frequencies towards either the European or the Vietnamese sampling sites. This suggests that, despite reduced coverage, introduced by sampling in the data set, the scored insertion polymorphisms are reliable. In addition, our method of analysis is likely to be conservative: indeed, the BAYESCAN outliers were selected for their consistency with a significant  $F_{CT}$  between European temperate and



**FIGURE 3** Insertion frequencies in Europe and Vietnam for 92 outlier loci. Bars represent the median value from the three reads sampling replicates, and dots represent the values from the other replicates (if outlier(s) found in replicates). Colours correspond to each of the five TE families



**FIGURE 4** Comparison of outlier frequencies with randomly selected loci. Insertion frequencies of 92 randomly chosen loci among those with the same minimum insertion frequency ( $\geq 20$  individuals) as outliers compared with 92 outlier loci. The random loci were taken from the first replicate (M1), and values for outliers are median values obtained from the three replicates. Nonoverlapping notches indicate a significant difference between the true medians (dark horizontal bars)

Vietnamese tropical populations, which avoids retaining outliers that we were not looking for, for example those due to a population-specific event.

Interestingly, we found significantly more outlier loci with a high frequency in Europe and low frequency in Vietnam. This was unexpected under our initial assumptions as follows: a favoured allele selected in one or another environment has a priori no reason to be more often associated with the presence or the absence of a TE insertion at linked sites. A possible explanation is that the majority of the sequenced TE insertions segregate at low frequencies (approximately 10% of all individuals). When considering the linked region of one polymorphic TE insertion, if a favourable mutation appears in an individual in which the insertion is absent, the increase in frequency of this “absence” haplotype will thus, most of the time, have a modest effect on the genetic differentiation at this marker because it is already segregating at high frequency. In contrast, if a favourable mutation appears in a TE “presence” haplotype, the increase in frequency of the linked TE insertion would lead to high differentiation ( $F_{CT}$ ). In absence of an alternative explanation, our outlier loci could thus indicate in which subset of populations the adaptive mutation occurred and in the present case this would have occurred more frequently in the temperate populations. Additionally, the observation of such a low global insertion frequency is expected to reduce the risk of false-positive outliers due to negative selection acting on potentially deleterious polymorphic insertions.

Two scenarios that are not mutually exclusive could be invoked in the light of our data. A simple case would be a direct adaptive evolution in European invasive populations that originated from tropical regions of the native area. A second hypothesis could be

that invasive temperate populations came from the northernmost territories of the native area such as northern China or Japan where *Ae. albopictus* populations are already cold-adapted. It would be thus interesting to know whether the observed signature of selection results from more “ancient” adaptations in the native area or if it originates from more recent fine tuning of cold-related traits in the invasive areas. A recent study (Porretta, Mastrantonio, Bellini, Somboon, & Urbanelli, 2012) using new variable COI mtDNA sequences and historical species range modelling suggested that northern territories of the native area of *Ae. albopictus* would be the latest to have been colonized after a range expansion from southern refugia after the last glacial approximately 21,000 years ago. The authors suggested that *Ae. albopictus* may have followed the human populations during their expansion from south to north in this area that began approximately 15,000 years ago. Thus, regardless of the origin of the invasive individuals sampled in Europe, it is likely that they are representatives of populations that had recently undergone a shift of selective pressure from tropical to temperate climatic conditions. This could explain why so many outliers are associated with high insertion frequency in Europe and why some candidate genes in the diapause pathway are found in the neighbourhood of some of these outliers. An easy method to distinguish between these possibilities would be to search if the same outlier insertions are present in several temperate populations from the native area.

We were able to assign a unique position for 21 of the outlier loci on the *Ae. albopictus* genome. Sixteen outliers were found inside or in the close vicinity of annotated genes, which may allow speculation on potential targets of selection. These genes encompass functions related to cell structure and organization, lipid metabolism or

signal transduction. A main challenge faced by temperate populations is overwintering (Denlinger & Armbruster, 2016; Hawley, 1988; Mori, Oda, & Wada, 1981; Takumi et al., 2009). Previous studies have shown that the cell cycle and lipid metabolism are specifically solicited at multiple stage of the diapause preparation and maintenance (Huang, Poelchau, & Armbruster, 2015; Poelchau, Reynolds, Elsik, Denlinger, & Armbruster, 2013b; Poelchau et al., 2013a; Urbanski et al., 2010) to allow temperate populations to go through winter within cold-resistant eggs. Although the genes found in our genome scan have not been functionally associated with diapause, two of them *laf1* (AALF000670) and a lipophorin-coding (AALF004790) are a differentially expressed between induced and noninduced samples during diapause preparation (Poelchau et al., 2013a) and are located closer than expected by chance from two of our outlier loci. These genes are both involved in lipid metabolism and could be thus strong candidates for adaptive evolution. Another notable candidate is AALF012643, a hemolymphatic carrier of the juvenile hormone (JH). JH appears to be critical in the maintenance of the diapause in *Ae. albopictus* but its exact function remains to be elucidated (Poelchau et al., 2013b). Although appropriate caution should be taken regarding the sole candidate status of these genes, it is worth mentioning that the diapause pathway has already been shown to benefit from rapid adjustments due to local adaptation in *Ae. albopictus*. For instance, Urbanski et al. (2012) showed that invasive American populations originating from Japan have rapidly evolved a new adaptive clinal response to diapause induction, independent from that observed in the native area. Thus, adaption in the temperate regions may have led to several selective sweeps on gene or regulatory sequences involved in this critical pathway, allowing the settlement of the mosquito in new temperate areas. Further experiments, including fine-scale study of the genetic diversity of these candidates among populations, are needed to assess their potential implication in the adaptation of *Ae. albopictus* towards temperate environments. Specifically, targeted resequencing of the candidate regions, including outliers, genes and their flanking regions across several individuals and populations should help to determine evidence of selective sweep, the precise extent and location of such events, and eventually designate the causative selected mutations.

It is also important to note that the results presented here are only restricted to a subset of the Asian tiger mosquito populations located in temperate and tropical environments. Thus, it is probable that some of the outliers detected could be specific to this particular comparison and do not reflect the global pattern of differentiation between tropical and temperate populations. Research on the same outliers between other tropical and temperate populations from native and non-native areas would be extremely valuable to extrapolate our results at a larger scale and refute possible false positive. Should the same outlier insertions be found at high frequencies in temperate locations—such as in USA, Japan or China—extended investigations about the origin of invasive populations would help clarify if those similarities are due to an ancestral sweep or parallel sweeps that occurred independently in several populations. This study already provides a set of functional primers for some

candidate loci that could be directly used to answer this question in *Ae. albopictus* DNA samples.

As with every novel method, our study may be susceptible to unforeseen or underrated bias; although we attempted to remove these issues (such as controlling the relationship between loci coverage and insertion frequency), we identified several points that must be taken into account for interpretation of the results. First, unlike random SNPs, TE insertions can be more often potential targets of purifying selection, which can sometimes mimic the diversity pattern of a selective sweep regarding the reduction in the genetic diversity (Charlesworth, Morgan, & Charlesworth, 1993; Stephan, 2010). However, as reported earlier, insertions are found at low frequencies and negative selection for the “presence” of TE is likely to produce a slight change in the allele frequency spectrum. Another issue may be the uncertainty related to the null model used in the genome scan; even if we did not explicitly evaluate the fit of our data to a specific population model, we did not find evidence to reject an island model such as that implemented in *BAYESCAN*. This software is also the best suited to handle our dominant data but is indeed restricted to mainly detect recent, strong and monogenic positive selection (Narum & Hess, 2011; Pérez-Figueroa, García-Pereira, Saura, Rolán-Alvarez, & Caballero, 2010; de Villemereuil et al., 2014); additionally, it should be insightful to compare our results with more diverse genome-scan models and we would like to emphasize that our results were produced under the specific hypothesis of the model used.

Here, we report the first information supporting adaptive evolution at the molecular level in the Asian tiger mosquito. Progress in the annotation of published genomes and the looming availability of supplementary genomic resources will allow the most gain from these results. We hope that this work will contribute to unravelling the implications of adaptive processes during the invasion of disease vectors.

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downloaded from [http://d-maps.com/carte.php?num\\_car=4719&lang=en](http://d-maps.com/carte.php?num_car=4719&lang=en) and [http://d-maps.com/carte.php?num\\_car=708&lang=en](http://d-maps.com/carte.php?num_car=708&lang=en). We thank the anonymous referees who contributed to the manuscript revision.

## DATA ACCESSIBILITY

Paired-end raw sequences are available through SRA at NCBI under SRP070185 (Bioproject PRJNA312147). Final presence/absence matrices (including replicates) are available at Dryad (<https://doi.org/10.5061/dryad.9p925>) at <https://doi.org/datadryad.org/review?doi=10.5061/dryad.9p925>.

## AUTHOR CONTRIBUTIONS

C.G., C.V. and M.B. conceived of the experiments and conducted the analyses. C.G. and H.H. developed and performed the molecular experiments. G.M., C.V.M. and P.M. conducted the sampling in France and Vietnam. All authors contributed to the final version of the manuscript.

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